(12)

EUROPEAN PATENT APPLICATION

- (43) Date of publication: 21.09.2005 Bulletin 2005/38
- (51) Int CI.7: C07C 233/00, C07C 237/00
- (21) Application number: 04006461.0
- (22) Date of filing: 18.03.2004
- (84) Designated Contracting States:
 AT BE BG CH CY C 2 DE DK EE SFIFR GB GR
 HU IE IT LI LLU MC NL PL PT RO SE SI SK TR
 Designated Extension States:
 AL LT LY MK
- (71) Applicant: Revotar Biopharmaceuticals AG 16761 Henningsdorf (DE)
- (72) Inventors:
 Kranich, Remo, Dr.
 13503 Berlin (DE)

- Aydt, Ewald Mirko, Dr. 10247 Berlin (DE)
- (74) Representative: Huhn, Michael et al Isenbruck Bösl Hörschler Wichmann Huhn Patentanwälle Theodor-Heuss-Anlage 12 68165 Mannheim (DE)
- (54) Non-glycosylated/-glycosidic/-peptidic small molecule selectin inhibitors for the treament of inflammatory disorders
- (57) The compounds having the general structure of formula (I)

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with R4 being H, CH3, CH2CH3

wherein the symbols, indices and substituents have the following meaning if R2=OH and R3=H then R1=H, CN, NO2, CF3, F, CI, Br, I, CH3 or

1, CH_3 of if $\text{R}^3 = \text{H}$ then $\text{R}^1 = \text{H}$, CN, NO_2 , CF_3 , F, CI, BI, I, CH_3 , Et, n-Pr, i-Pr, n-Bu, i-Bu, t-Bu, phenyl, thienyl, furyl, thiazolyl or

if R3=OH and R1=H then R2=H, CN, NO₂, CF₃, F, Cl, Br, I, CH₃, Et, n-Pr, i-Pr, n-Bu, I-Bu, phenyl, thienyl, furyl, thiazolyl

then X is e.g.

and Y being

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and the pharmaceutically acceptable salts, esters or amides and prodrugs of the above identified compounds of formula (f). The compounds are applied to modulate the in-vitro and in-vivo binding processes mediated by E., P- or L-selectin binding.

Description

[0001] Compounds and methods are provided, which modulate the in-vitro and in-vivo binding processes mediated by E-selectin, P-selectin or L-selectin binding. More specifically, selectin inhibitors and their use are described, wherein the selectin inhibitors that inhibit the selection-mediated function comprise polyhydroxy obenyl subunits.

[0002] When a tissue has been invaded by a microorganism/is intected or has been damaged, the inflammatory process directs leukocytes and other immune system components to the site of infection or injury. Within this process. leukocytes (white blood cells) play a major role in the engultment or digestion of microorganisms. Thus, the recruitment of leukocytes to injected or gamaged tissue is critical for mounting an effective immune defense. Generally, white blood cells are found circulating through the bloodstream. To migrate from the blood stream into the affected tissue, the white blood cells must be able to recognize the invaded or damaged lissue and be able to bind to the wall of the capillary (endothelium) near the affected tissue and diffuse through the capitlary into the affected tissue. Therefore, leukocytes have to roll and adhere to the endothelial cell wall. This cell adhesion event is one of the most important aspects of the inflammatory response. The tirst steps of this cell adhesion are mediated by members of the selectin family. The selectin tamily of vascular adhesion molecules is comprised of three structurally related calcium-dependent carbo hydrate binding cell surface proteins, P. L. and E-selectin, E-selectin is expressed only on inflamed endothelium, P. selectin is expressed on inflamed endothelium as well as on platelets, and has much structural similarity to E-selectin, L-selectin is expressed on leukocytes and also has much structure similarity to P- and E-selectins. The selectins are transmembrane proteins and are composed of an amino terminal lectin domain, an epidermal growth factor (EGF)-like domain, a variable number of complement receptor-related repeats, a hydrophobic domain spanning region and a cytoplasmic domain. The binding interactions which lead to the adhesion of the leukocytes appear to be mediated by contact of the lectin domain of the selectins and various carbohydrate ligands on the surface of the leukocytes. All three selectins can bind to the carbohydrate sialyl Lewis*, a glycosyl molety present on the surface of most/certain leukocytes. In case of P- and L-selectin, a distinct protein liband has been described [R.P. McEver; R.D. Cummings; J.Clin.Invest.; 1997; 100; 485-492), the so-called PSGL-1 (P-selectin glycoprotein ligand-1), which contributes to a high affinity selectin binding by its sLex moiety as well as by parts of its peptide components, sulphated tyrosine (R.P. McEver; Ernst Schering Res. Found, Workshop; 2004; 44: 137-147), PSGL-1 is one of the most important selecting ligands binding with highest affinity to P-selectin, but it also binds to E-and L-selectin. It is expressed as homodimenc sialomucin on leukocytes and platelets where it is upregulated during inflammation.

[0003] In contrast to her low beal expression, E- and P-selectin expression is also upregulated during inflammation, leading to a substantial recruitment of leukocytes into the inflamed tissue. Although selectin-mediated cell adhesion is required for lighting infection, there are situations in which such cell adhesion is undestrable or excessive, resulting in tissue damage instead of repair. In the case of many acute as well as chronic inflammatory disorders (e.g., asthmat, chronic obstructive pulmonary disease (COPD), positissis, etc.) an association between infiltration of activated feukocytes into the lissue simultaneously with a marked elevation of tissue expression of corresponding adhesion molecules, particularly E- and P-selectin, has been demonstrated. Such abnormal white blood cell infiltration, caused by vertice selectin expression may also play a role in transplant and graft rejection. Also the process of blood clotting is further promoted by leukocyte-leukocyte and leukocyte-platelet brinding, which occurs because leukocyte possess as welluselectin as its corresponding ligand PSGL-1 and can thus interact with themselves, via PSGL-1, and they can also brind to platelets which carry P-selectin.

[0004] D. Bock et al., New Drugs. p. 28 to 30 (28.04.2003) describe the role of selectin antagonists, especially bimoslamose, as inhibitor of selectins being an inflammation target;

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[0005] As additional targets MIF (Macrophage Migration Inhibitory Factor) and JAM-1 (Junctional Adhesion Molecule 1) are mentioned.

[0006] EP 0 758 243 B1 describes Mannopyranosyloxy-phenyl-benzoic acid or similar acids as components in a medicine for treating or preventing diseases, characterized by the binding of E-, P- and/or L-selectin to a sialyl-Lewis x (sLe*) or sialyl-Lewis at (sLe*) presented on a cell surface through the mibition of such binding, Beades inflammatory diseases, other diseases like septic shock, psoriasis, feurnatoid anthritis, reperfusion injury and cancer are mentioned. [0007] From EP 0 840 668 B1 compounds like bimosamose are known, e.g. 1, 16-Bis-Ji-carboxymethylphenyl)-4-(2-or-D-mannopyranosyloxylphenyllhexan and derivatives thereof including heptan, butan and pentan derivatives. Again, their use for inhibiting the selectin binding to sLe* or sLe* are mentioned.

Q (0008) A different method for inhibiting the binding between a first cell having a selectin and a second cell having a ligand for selectin in vitro is described in EP 0 902 881 B1 in permitting a light domposition to interact with the first cell. A proportion of the lights has an attached saccharide like fuccoligosaccharides or lactose, a different proportion has, e.g., a second attached acidic monosaccharide, further proportions are crosslinked and a proportion without attached saccharides have an acidic group being negatively charged at neutral pH. These lipid compositions are used in local attentions in the adherence of leukocytes or cancer cells to vascular endothellum, platelets or lymphatic tissue.

[0009] EP 1 288 222 A1 describes compounds with affinity to P-selectin being peptides with a specific amino acid sequence comprising, e.g., valine, tryptophan, aspartic acid, and D-ort-glutamic acid. Medicaments comprising these peptides are said as inhibiting leukocyte binding to platelets and/or endothelial cells. Similar peptides are known from WO-A 03/020753.

[0010] The disalicylates and disalicylate-based C-glycosides of WO-A 99/29708 act as selectin-ligand structural mimetics in medicaments. They may lack the sialic acid and/or fucose of the natural selectin ligand being sLe^x. A typical compound is

[0011] The 5-membered heterocycles of WO-A 00/33836 exhibit Inhibitory activity against the selectins; typical structures are

with substituents like a moiety containing a terminal carboxylic acid group as a non-N substituent, a hydrophobic moiety of such as a functionalized alkyl chain as a non-N substituent and a functionalized aryl group which also may be an N-substituent.

[0012] MO: A 01/89531 provides methods for identifying agents which interact with (activate or inhibit) the crystal and three-dimensional structures of a) the lection and EGF-like (LE) domains of P-selectin, of b) P-selectin LE each complexed with sLe*, and c) P-selectin LE complexed with a functional PSGL-1 peptide modified by both trosains explaint and staff size.

[0013] The compounds for modulating in vitro and in vivo processes mediated by selectin binding according to WO-A 03/097658 are benzyl amino sulfonic acids linked to carbohydrate or glycomimetic. A typical compound would be

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C.C.M.Appeldoom et al. in JPC Papers in Press, published January 13, 2003, with the title "Optimization of a Pselectin briding peptide" report on peptides containing a GNu-Try-14-Asp-Val consensus motif to which gallic add or 1,3,5-ben-zenetricarboxylic acid at the N-terminus have been added as substituents, C-terminus modifications had little to no influence on the affility of the core peptide. The two trypical substituents are

is a typical peptide rest.

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[0014] In such a very complex pharmaceulical field, there is a strong medical need in the art for indentifying further inhibitors of selectin-dendated function. e.g. of selectin-dependent ceil adhesion, and for the development of methods employing such compounds to inhibit conditions associated with excessive selectin activity. Most of the available pharmaceulical therapies e.g. of psoriasis or asthma, which are available on the market comprise mostly conficosteroids or NSAIDs (non steroidal anti-inflammatory drugs) which have several serous drawbacks/side effects, and target different steps of the inflammatory cascade.

[0015] Object of the invention is to provide new compounds, especially non-glycosylate/dlycosidic relations in molecules, which are able to inhibit selection and which have less negative side effects during their in-vitro or investigation. Furthermore, the said molecules should show a good bioavailability when applied as an active ingredient in a medicine.

[0016] The present invention provides compounds having the general structure of formula (I).

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wherein the symbols, indices and substituents have the following meaning

if R^2 =OH and R^3 =H then R^1 =H, CN, NO₂, CF₃, F, Cl, Br, I, CH₃ or if R^3 =OH and R^2 =H then R^1 =H, CN, NO₂, CF₃, F, Cl, Br, I, CH₃, EI, n-Pr, i-Pr, n-Bu, i-Bu, i-Bu, phenyl, thienyl, furyl,

If R3=OH and R1=H then R2=H, CN, NO₂, CF₃, F, CI, Br, I, CH₃, E1, n-Pr, i-Pr, n-Bu, t-Bu, phenyl, thienyl, furyl, thiazolyl -X- =

(a)

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with -E- being -NH-, -(CH₂-)_kNH-; -G- being -(NH-)_m and g = 0,1; h = 1, 2, 3; k = 1, 2, 3; m = 0,1; n = an integer from 1 to 8

(b)

No.

with R4 being H, CH3, CH2CH3

(c)

with $\rm R^5$ being H, NO $_2$, CF $_3$, F, CI, Br, I, CN, CH $_3$ and -K- being -S- or -O-

(d)

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with p being an integer from 2 to 8

55 (e)

with q being an integer from 1 to 9 and r being an integer from 1 to 3

with -T1-, -T2-being -E-, -K- or -(N-alkyl)--Y =

with -V- being -(NH-)s- with s being 0 or 1,

[0017] R^6 being CO_2H , $CO_2Alklyl$, CO_2Alyl , CO_2NH_2 , $CO_2Aralkyl$, SO_3H , SO_2NH_2 , $PO(OH)_2$, 1-H-letrazolyl-, CHO, $COCH_3$, CH_2OH , NH_2 , NICAlkyl, NICAlkyl

-W-R⁶

5 [0018] Preferred compounds are those of formulas (A) and (B)

wherein -Y is (Q=CH, N)

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 $\label{eq:with R10-being CO2H, CO2elkyl, CO2aryl, CO2NH2, CO2aralkyl, CH2SO3H, CH2SO3H, CH2FO(OH)2, 1-H-letrazolyl, CH0, COCH3, CH2OH, CH2NH2, CH2Nlalkyl, CH2N(alkyl)alkyl', CH2OCH3, CH2SH, CH2NLalkyl, CH2NLalkyl) alkyl', CH2OCH3, CH2SH, CH2NLalkyl, CH2NLa$

 R^{11} being CO_2H , $CO_2\text{alkyl}$, $CO_2\text{aryl}$, $CO_2\text{NH}_2$, $CO_2\text{aralkyl}$, SO_3H , $SO_2\text{NH}_2$, $PO(OH)_2$, 1-H-tetrazolyl, CHO, COCH_3, OH, NH2, NHalkyl, N(alkyl)alkyl', OCH_3, SH

[0019] Further particularly preferred compounds include the following formulas (C) and (D)

wherein Y is like relating to formulas (A) and (B) and X is:

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[0020] Also part of the invention are the pharmaceutically acceptable salts, esters or amides and prodrugs of the above-identified compounds of formula (I).

The present invention also provides pharmaceutical compositions comprising at least one compound of the formula (1) and a pharmaceutically acceptable carrier which is useful in a medicine. These compounds present the ability of inhibiting cell adhesion and inhibit selectin-[as well as PSGL-1-mediated] binding. The compounds have the ability to inhibit the interaction of selectins with sLe* / sLe* and also the interaction between selectins and tyrosinesullate resi-

In a preferred variant the invention provides pharmacoutical compositions comprising at least one compound of formula (A) or formula (B) or formula (C) or formula (D) and a pharmacoutically acceptable carrier. The present invention further provides a method of inhibiting the binding of P-selectin, t-selectin or E-selectin to stee* or stee* and tyrosinesulfate residues comprising the step of administering to a patient an effective amount of at least one compound having the structure of formula (1) to inhibit the binding of P,E- or t_selectin to stee* or stee* and tyrosinesulfate.

[0021] It has been found that compounds having the formula (f) shown above act to inhibit E, P, or L-select P or 10 or 1

The term "aryl" shall mean carbocyclic and heterocyclic aromatic groups including, but not limited to, phenyl, 1-naphthyl, 2-naphthyl, fluorenyl, (1,2)-dihydronaphthyl, indenyl, indanyl, thienyl, benzothienyl, thienopyridyl and the like.

The term "aralky" (also called arylalky) shall mean an any group appended to an alkyl group including, but not limited to, benzyl, 1-naphthymethyl, 2-naphthymethyl, fluorobenzyl, chlorobenzyl, bromobenzyl, iodobenzyl, alkoxykenzyl (wherein "alkoxy" means methoxy, ethoxy, isopropoxy, n-butoxy, sec-butoxy, tert-butoxy an the like), hydroxybenzyl, aminobenzyl, nitrobenzyl, guanidinobenzyl, fluorenylmethyl, phenylmethyl(benzyl), 1-phenylethyl, 2-phenylethyl, 1-naphthylethyl and the like.

[0022] The term "pharmaceutically acceptable salts, esters, amides and prodrugs" as used herein refers to those carboylate salts, amino acid addition salts, esters, amides and prodrugs of the compounds of the present invention which are, within the scope of sound medical judgement, suitable for use in contact with tissues of patients without under toxicity, irritation, altergic response and the like, commensurate with a reasonable benefit/risk ratio, and effective

for their intended use, as well as the zwitterionic forms, where possible, of the compounds of the present invention. The term "salts" refers to the relatively non-toxic, inorganic and organic acid addition salts of the compounds of the present invention. These salts can be prepared in situ during the final isolation and purification of the compounds or by separately reacting the purified compounds in its tree form with a suitable inorganic or organic acid or base and isolating the salt thus formed. Representative salts of the compounds of the present invention include the hydrobromide. hydrochloride, sullate, bisulfate, nitrale, acetate, oxalate, valerate, palmitate, stearate, laurate, borate, benzoate, lactate, phosphate, tosylate, citrate, maleate, fumarate, succinate, tartrate, naphthylate, mesylate, glucoheptonate, lactiobionate, laurylsulphonate salts and the like. These may include cations based on the alkali and alkalineearth metals, such as sodium, lithium, potassium, calcium, magnesium and the like, as well as non-toxic ammonium, quaternary ammonium and amine cations including, but not limited to, ammonium, tetramethylammonium, tetraethylammonium, methylamine, dimethylamine, trimethylamine, triethylamine, ethylamine, and the like.

Examples of the pharmaceutically acceptable, non-toxic esters of the compounds of this invention include C1, C2, C3, C4. C5 and C6 alkyl esters wherein the alkyl group is a straight or branched chain. Acceptable esters also include C5. C6 and C7 cycloalkyl esters as well arylalkyl esters such as, but not limited to benzyl. C1, C2, C3, C4, C5 and C6 alkyl ester are preferred. Esters of the compounds of the present invention may be prepared according to conventional methods.

[0023] Examples of pharmaceutically acceptable, non-toxic amides of compounds of this invention include amides derived from ammonia, primary C₁, C₂, C₃, C₄, C₅ and C₆ alkyl amines and secondary C₁, C₂, C₃, C₄, C₅ and C₆ dialkyl amines wherein the alkyl groups are straight or branched chains. In the case of secondary amines the amine may also be in the form of a 5 or 6 membered heterocycle containing one nitrogen atom. Amides derived from ammonia, C_1 , C_2 and C3 alkyl primary amides and C1 to C2 dialkyl secondary amides are preferred. Amides of the compounds of the present invention may be prepared according to conventional methods.

[0024] The term "prodrug" refers to one or more compounds that are rapidly transformed in vitro and from a nonactive to active state in vivo to yield to the parent compound of the above formula (1), for example by hydrolysis in blood

or in vivo metabolism.

The present invention also provides torpharmaceutically active compositions that contain the compounds of the present invention. It is also contemplated that pharmaceutically active compositions may contain a compound of the present invention or other compounds that inhibit or compete with E-selectin or P-selectin or L-selectin binding.

Pharmaceutically active compositions of the present invention comprise a physiological carrier and a compound of formula (I). The pharmaceutical compositions of the present invention may include one or more of the compounds having the above structure (I) formulated together with one or more non-toxic, physiologically acceptable carriers, adjuvants or vehicles, which are collectively referred to herein as carriers, for parenteral injection, for oral administration in solid or liquid form, for rectal or topical administration and the like.

The compositions can be administered to humans and animals either orally, rectally, parenterally (intravenously, intramuscularly, or subcutaneously), intracisternally, intravaginally, interperitoneally, locally (powders, ointments or drops), or as a buccal or by inhalation (nebulized, or as nasal sprays).

Compositions suitable for parenteral injection may comprise physiologically acceptable sterile aqueous or nonaqueous solutions, dispersion, suspensions or emulsions and sterile powders for reconstitution into sterile injectable solution or dispersion. Examples of suitable aqueous and nonaqueous carriers, diluents, solvents or vehicles include water, ethanol, polyol, (propylene glycol, polyethylene glycol, glycerol and the like), suitable mixtures thereof, vegetable oils (such as olive or canola oil) and injectable organic esters such as ethyl oleate. Proper fluidity can be maintained, for examples, by the use of a coating such as lecithin, by the maintenance of the required particle size in the case of dispersions and by the use of surfactants.

These compositions may also contain adjuvants such as preserving, wetting, emulsifying, and dispersing agents. Prevention of the actions of microorganisms can be ensured by various antibacterial and antifungal agents, for examples, parabens, chlorobutanol, phenol, sorbic acid, and the like. It may also be desirable to include isotonic agents, for examples sugars, sodium chloride and the like. Prolonged absorption of the injectable pharmaceutical form can be brought about by the use of agents delaying absorption, for examples aluminium monostearate and gelatin.

If desired, and for more effective distribution, the compounds can be incorporated into slow or timed release or targeted delivery systems such as polymer matrices, liposomes, and microspheres. They may be sterilized, for example, by tiltration through a bacteria retaining tilter, or by incorporating sterilizing agents in the form of sterile water, or some other sterile injectable medium immediately before use.

Solid dosage forms for oral administration include capsules, tablets, pills, powders and granules. In such solid dosage forms, the active compound or a prodrug ester is admixed with at least one inert customary excipient (or carrier) such as sodium citrate or dicalcium phosphate or (i) fillers or extenders, as for example, starches, lactose, sucrose, glucose, mannitol and silicic acid, (ii) binders, as tor example, carboxymethylcellulose, alginates, gelatine, polyvinylpyrrolidone, sucrose and acacia. (iii) humectants, as for example, glycerol, (div disintegrating agents, as for example, agar-agar, calcium carbonate, potato or tapioca starch, aliginic acid, certain complex silicates and sodium carbonate,

(v) solution relarders, as for examples, paraflin, (vv) absorption accelerators, as for example, quaternary ammonium compounds, (vii) wetting agents, as for examples, cetyl alcohol and glycerol monstearate, (viii) adsorbents, as for 'example, kaolin and bentonite, and (ix) lubricants, as for example, talc, calcium stearate, magnesium stearate, solid polyethylene glycols, sodium lauryl sulfate and mixtures thereof. In the case of capsules, tablets, and pills, the dosage forms may also comprise buffering agents.

Solid compositions of a similar type may also be employed as titlers in soft and hard-titled gelatine capsules using excipients as factore or milk sugars as well as high molecular polyethylene objects and the like.

Solid dosage forms such as tablets, dragées, capsules, pills and granules can be prepared with coatings and shelle, such as enteric coatings and others well known in the art. They may contain opacitying agents, and can also be of such compositions that they release the active compound or compounds in a certain part of the intestinal tract in a delayed manner. Examples of embedding compositions that can be used are polymeric substances and waste.

The active compounds can also be in microencapsulated form, if appropriate, with one or more of the abovementioned excipients

Liquid dosage forms for oral administration include pharmaceurically acceptable emulsions, solutions, suspensions, syrus and elixirs. In addition to the active compounds, the liquid dosage forms may contain inert disturts commonly used in the art such as water or other solvents, solubilizing agents and emulsifiers, as to example, ethyl alcohol, isopropylatchol, ethyl carbonate, ethyl alcohol, benzyl benzoate, propylene glycol, 1,3-butylene glycol, dimethylloramanide, oils, in particular, cottonseed oil, groundrul oil, corn germ oil, olive oil, cannola oil, caster of and sesame seed oil, glycordnot, oliven oil, cannola oil, caster of and sesame seed oil, glycordnot, ethyl or other or mixtures of these substances, and the like.

Besides such inert diluents, the compositions can also include adjuvants, such as wetting agents, emulsifying and suspending agents, sweeting, flavouring and perfuming agents.

Suspensions, in addition to the active compounds, may contain suspending agents, for example, ethoxylated isostearyl alcohols, polyoxyethylene sorbitol and sorbian esters, microcrystalline cellulose, aluminium metahydroxide, bentonite, agar-agar, tragacanth or mixtures of these substances and the like.

Compositions for rectal administrations are preferably suppositories, which can be prepared by mixing the compounds of the present invention with suitable noniritating excipients or carriers such as cease butter, polyether glycol or a suppository wax, which are solid at ordinary temperatures but liquid at body temperature and therefore melt in the rectal or varianja cavity and release the active component.

Dosage forms for topical administration of a compound of this invention include ointments, powder, sprays and inhalants.

The active component is admixed under sterile conditions with a physiologically acceptable carrier and any neededge-servatives, buffers or propellants as rhay be required. Ophthalmic formulations, eye ointments, suspensions, powder and solutions are also contemplated as being within the scope of this invention.

The compounds of the present invention can also be administrated in the form of [posomes. As is known in the art, lipsoemes are generally derived from phospholipids or other injied substances. Liposemes are formed by mon or multilamellar hydrated liquid crystals that are dispersed in an aqueous medium. Any non-toxic, physiologically acceptable metabolized lipid capable of forming iposomes can be used. The present compositions in liposome form can contain, in addition to the selectin binding inhibitors of the present invention, stabilizers, preservatives, excipients and the like. The preferred lipids are the phospholipids and the phosphatidylcholines (lecithins), both natural and synthetic. Methods to form [posomes are well known in the art.

Actual desage levels of active ingredient in the composition of the present invention may be varied so as to obtain an amount of active ingredient that is effective to obtain the desired therapeutic response for a particular composition and method of administration. The selected desage level, therefore, depends on the desired therapeutic effect, on the route of administration, or the desired duration of treatment and other factors.

The total daily dosage of the compounds on this Invention administered to a host in single or divided doses may be in the range up to 5 on mg per kilogram of body weight. Dosage unit compositions may contain such submultiples thereof as may be used to make up the daily dosage, it will be understood, however, that the specific dose level for any particular patient, whether human or other animal, will depend upon a variety of lactors including the body weight, general health, sex diet, time and route of administration, rates of absorption and excretion, combination with other drugs and the severity of the particular disease being retained.

In particular, the compounds of the present invention may be used to treat a variety of diseases relating to inlammation and cell-cell recognition and adhesion. For example, the compounds of the present invention may be administrated to a patient to treat COPD, acute respiratory distress syndrome (ARDS), Crohn's disease, septic shock, chronic inflammatiory diseases such as psoriasts, atopic dematitis, and rheumatioid arthritis, and repertusion tissue injury that occurs following heart attacks, storkes, atheroselerosis, and organ transplants, traumatic shock, multi-organ failure, autoimmune diseases like multiple sclerosis, asthma and inflammatory bowel disease. In each case, an effective amount of the compounds of the present invention is administered either alone or as part of a pharmaceutically active

composition to a patient in need of such treatment. It is also recognized that a combination of the compounds may be administered to a patient in need of such administration. The compounds of the present invention may also be administrated to treat other diseases that are associated with cell-cell adhesion. As the present compounds inhibit the binding of E-selectin or P-selectin or L-selectin, any disease that is related to this interaction may potentially be treated by the inhibition of this binding interaction.

In addition to being found on some white blood cells, s.Le^a is found on various cancer cells, including lung and colon cancer cells. It has been suggested that cell adhesion involving s.Le^a may be involved in the metastasis of certain cancers and inhibitors of s.Le^a binding might be useful in treatment of some forms of cancer.

[0025] Many of the compounds of the present invention may be synthesized according to the following general synthetic schemes.

In SCHEME 1 an amino acid of type (1) is reacted to the corresponding methyl ester (2) under heating with actide methanic Ester (2) is reacted with a trimethoxy-phenyl-allytic acid under state-of-the-ant conditions (i.e. N°-G-dimethylaminopropyl)-N-ethyl carbodirinide (EDC), triethylamine and 4-dimethylaminoprydine (DMAP) in a chlorinated solvent) to the aminoid (3). Alternatively dispopropyl carbodirinide (D(C) and hydroxybenzotitazele (HOBI) may be used for this reaction step. Amide (3) is converted to acid (4) by treating it with an excess of boron tribromide at -78°C up to it in a halogenated solvent followed by aqueous workup. However, in some cases it may be necessary to hydrotize the ester (3) with it.e. aqueous lithium hydroxide in acctonitritle before treating it with boron tribromide. The synthesis sequence shown in SCHEME 1 leading to compounds like (4) is not only reduced to the Y-H building blocks like (1) but may be generally applied to all other Y-H type building blocks leading to compounds of type (A) and (B) as shown in the paragraph before.

SCHEME 2

In SCHEME 2 a carboxy substituted thiophene like (5) is reacted to the corresponding ethyl ester (6) under heating in acidic ethanol. Ester (6) is tornimated with N-bromosuccinimide in anhydrous chlorotorn and glacial aceitic acid to give (7) which is further reacted with 2-Amino-benzeneboronic acid under a state-of-the-art Suzuki transformation (i. e. Tetrakis(friphenyiphosphine)-palladium, aqueous sodium carbonates, ethanol, toluene) to the biapy (8). Biapyl (8) is reacted with a timenthosy-phenyi-alkylic acid, EDC, triethylamine and DMAP in a chlorinated solvent to the amide. Alternatively DIC and HOBI may be used for this reaction step. Amide (9) is converted to acid (10) by treating it with an excess of boron tribromide at -78°C up to it in a halogenated solvent followed by aqueous workup. However, in some cases if may be necessary to hydrolize the ester (9) with i.e. aqueous lithium hydroxide in acetonlitrile before treating it with boron tribromide.

SCHEME 3

In SCHEME 3 4-Amino-butyric acid hydrochloride (11) is reacted at rt with an trimethoxybenzoic acid under basic conditions (triothylamine in anhydrous dichloromethane) to the amide (12) which is further hydrolized with aqueous lithium hydroxide in acetonitrie to give a building block like (13).

SCHEME 4

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In SCHEME 4 an acid like (13) is reacted with an aniline of general type (14) under state-of-the-art conditions (i.e. EDC, triethylamine and DMAP in a chbrinated solvent) to form the amide (15). Alternatively DIC and HOBI may be used for this reaction step. Amide (15) is converted to acid (16) by treating it with an excess of boron tribromide at 7,8°C up to it or in a halogenated solvent followed by aqueous workup.

In case that R⁶ and/or R⁸ contain carboxylic acid functionalities, those are protected as their corresponding methyl or ethyl esters before and hydrolized alterwards to release the carboxylic acid functionalities. The ester hydrolysis was done whether with LIOH in McCN or under treatment with BR₂ in followed by addition of water.

The synthesis sequence shown in SCHEME 4 leading to compounds like (16) is not only reduced to X-Y-H and Y-H building blocks like (14) but may be generally applied to all other X-Y-H and Y-H type building blocks leading to compounds of type (C) and (D) as shown in the paragraphs before.

SCHEME 5

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For compounds of type (23) containing a carboxylic acid functionality a synthesis sequence according to SCHEME 5 may be applied. An aniline of type (17) is reacted at a with Pulurenymethorycarbony of bindried (Fmoc-C) in 1.4 dio xane and aqueous sodium carbonate to the corresponding Fmoc-protected compound (18). Acid (18) is immobilized on solid support by esterification with 2-Chlorotrinjchhoride-polystyrene and Huenig's base in annytorous dichlorometh-ane and N.N-dimethyllormamide at 1 to give (19) which is threft retated with piperdine in N.N-dimethyllormamide. The resulting compound (20) is reacted then with acid (13), DIC and HOBI in N.N-dimethyllormamide at 1 to give itself with 1.1,1,3,3-li-lexalitycore-2-propanol in dichloromethane at 1 to cleave (22) from soid support. Following treatment of (22) with an excess of boron tribromide in dichloromethane at -78°C up to rt and aqueous workup provides the desired compound (23).

The present invention is furthermore illustrated by the following representative examples.

(3-[3-(2,3,4-Trihydroxy-phenyl)-propionylamino]-phenyl)-acetic acid (27)

[0026]

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SCHEME 6

[0027] Step 1: Dissolve (3-Amino-phenyl)-acetic acid ((24),700mg, 4-63mmd) in MeOH ((21mL) and add conc. subfuric acid (0.27mL, 5.06mmd). Stir her heaction mixture for 2d under reflux. Cooled mixture to 1 for memorature (ff), remove solvent under reduced pressure and prepurity the residue by flushing it over a short pad of silica gel using CIOAc. Remove solvent again and partitione the residue between EIOAc and saturated aqu. NaHCO₂ (1+1). Extracte the aqueous layer 3 times with EIOAc, washe the combined organic layers with brine and dried with Na₂SO₄. Remove solvent under reduced pressure and dry the residue without further purification in oil pump vacuum to obtain product (25) as a light yellow oil (708mg, 92%). "H NMR (400MHz, CDCI): 3.51 (s, 2 H); 3.67 (s, 3 H); 6.57 (dd, 1 H, J=78Hz), 7.8Hz, J₂= 1.8Hz); 6.60 (kr.4, 1, H, J=7.8Hz), 6.65 (kr.4, H, J=7.8Hz), 7.98 (Yr.1, H, J=7.8Hz),

[0028] Step 2: (The following reaction is done in an anhydrous N₂ atmosphere.) Dissolve EDC hydrochloride (187mg, 0.98mmol) and triethylamine (0.14mL, 1.00mmol) in anhydrous dichloromethane (3.5mL) and stir for 5min at rt. Added 3-(2.3,4-Trimethoxy-phenyl)-propionic acid (234mg, 0.97mmol) and DMAP (12mg, 0.10mmol) and stir for 10min. Added ester (25) (107mg, 0.65mmol) and stir for reaction solution overnight at rt.

Hydrolize the reaction solution with saturated aqu. NH_xCI followed by water, separate layers, extracte aqu. layer with dischioromethane ig times) and washe the combined organic layers with water and brine and dry with Na₂SO₄. Remove solvent under reduced pressure.

Purily crude product by preparative radial chromatography (silica gel 60PF, ELOACC/H1+1) to obtain product (26) as white solid (209mg, 83%), (K. C. Nicolaou; P. S. Baran, Y.-L. Zhong; K. Sugits; J. Am. Chem. Soc.; 2002; 124; 10; 2112;2220; 1+ NMRI (400MHz, CDClg); 2 sc2 (t, 2 H, J = 7.5Hz); 2.95 (t, 2 H, J = 7.5Hz); 3.58 (s, 2 H); 3.67 (s, 3 H); 3.92 (s, 3 H); 3.94 (s, 3 H); 3.91 (s, 3 H); 6.59 (d, 1 H, J = 8.6Hz); 6.96 (d, 1 H, J = 8.6Hz); 6.98 (br.d, 1 H, J = 7.8Hz); 7.92 (Yt, 1 H, J , 7.8Hz); 7.93 (br.d, 1 H, J = 7.8Hz); 7.92 (Yt, 1 H, J , 7.8Hz); 7.94 (br.d, 1 H, J = 7.8Hz); 7.94 (br.d

[0029] Step 3: (The following reaction is done in an anhydrous N₂ atmosphere.) Dissolve (26) (139mg, 0.36) in anhydrous dichloromethane (1.8mL), cooled to -78°C (acetone/ dy ice bath) and add slowly BBr₃ (0.5 1mL, 5.39mmol). Sit the reaction mixture for additional 30min at -78°C. Remove cooling bath and sit the reaction mixture for 3°D at 1 the Cool reaction mixture to 0°C, add slowly water (0.50mL) under vigorous stiring followed by dichloromethane (1.0mL) and methanol (2.0mL). Extract the mixture with EIOAc (3 times) and dry the combined organic layers with Na₂CO₂Remove solvent under reduced pressure and purify crude product by preparative RP HPLC (gradien, water/MeCN

95:5) to obtain (3,13-(2,3,4-Tirhydroxy-phenyl)-propionylaminol-phenyl)-acetic acid (27) as a white solid (68mg, 57%) IT. P Kogan: B. Dupré; H. Bui; K. L. McAbee; J. M. Kassir; I. L. Scott, X. Hu; P. Vanderslice; P. J. Beck; R. A. F. DixonJ. **Med. Chem.; 1998; 41; 1099-1111]. **H NMR (400MHz, CO₂OD); 2.67 (1, 2 H. J = 7,6Hz); 2.92 (1, 2 H. J = 7,6Hz); 3.61 (s. 2 H), 6.31 (d. 1 H. J = 8,3Hz); 5.50 (d. 1 H. J = 8,3Hz); 7.05 (brd, 1 H. J = 7,8Hz); 7.28 (YI, 1 H. J = 7,8Hz); 7.48 (brd, 1 H. J = 7,8Hz); 7.78 (brd, 1 H. J = 7,8Hz); 7.49 (brd, 1 H. J = 7,8Hz); 7.40 (brd, 1 H. J = 7,8Hz);

EXAMPLE 2

(5-{2-[2-(2.3,4-Trihydroxyphenyl)-acetylamino]-phenyl}-thiophen-2-yl]acetic acid (34)

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SCHEME 7

[9031] Step 1: Dissolve Thiophene-2yl-acetic acid (28) (2.44g, 17.1mmol) in ethanol (35mL) and add furning aguhydrochloric acid (few drops). Sift the reaction mixture for 19 ha 170 °C. Cool mixture to 11, remove solvent under reduced pressure and resolve the residue in EtOAc. Wash this organic tayer 3 times with 5% agu. Na₂CO₂ and extract the combined aqueous tayer 3 times with EtOAc. Wash the combined organic tayers with brine and dry with Na₂SO₄, Remove solvent under reduced pressure and dry the residue without further purification in oil pump vacuum to bind product (29) as a light brown oil (2.78g, 55%). [J. Kunes; V. Balsanek, M. Pour; V. Buchta; Collect. Czech. Chen. Commun. 2001. 65: 12; 1809-1830; 11 NIMR (400MHz, CDCI); 1.26 (1.9, 1.7, 1.1 Hz); 381 (5.2 H); 4.17 (6.2 H.

J = 7.1 Hz); 6.91-6.96 (m, 2 H); 7.20 (d, 1H, J = 4.8 Hz).

[0032] Step 2: (The following reaction is zone in an anthydrous N_aimosphere) Dissolve ester (29) (1.30g, 7.64mmol) in anthydrous chioroform (6.0mL) and glacial acetic acid (6.0mL), add N Bromosucchimided (1.39g, 7.79mmol) in portions and stir the mixture for 23h at rt. The mixture is diffused with an equal volume of water, the organic layer separated and washe with a 1M agu, NaOH, water, again with 1M agu, NaOH and water (2 times). Finally wash the organic layer with brine and only with Na₂OA, Remove solvent under reduced pressure. Purily crude product by preparative radial chromatography (silica gel 60PF, CyH/E(CAC 5+1) to obtain product (30) as an impured (according to NME; 20% sideproduct) orange liquid (1.61g, 55%) which is used without any further purification. [P. M. Jackson; C. J. Moogy: P. Sha; J. Chom. Soc. Perkin Trans. 1; 1990; 2009-2918; 1+1 NMR (400MHz, CDClg)* 1.26 (t. 3 H, J = 7.1 Hz), 3.73 (s. 21); 4.17 (g. 21, H, J = 7.1 Hz), 5.67 (d. 11, H, J = 3.5 Hz); 6.86 (d. 11, H, J = 3.5 Hz).

2 H): 4.17 (g, 2 H, J = 7.1 Hz); 6.67 (d, 1 H, J = 3.5 Hz); 6.88 (g, 1 H, J = 3.5 Hz).
[0033] Steps: (The tollowing reaction is done in an oxygentree N₂ aimosphere): ethanol (1.47mL). Tetrakis-(triphenylphosphine): palladium(0) (59.0mg, 2.5moR%) and agu. Na₂CO₃ (1.60g, 5.60mmol); presolved in 2.0mL H₂O) are subsequently added to discoved 2-Aminoberneehoronic acid (341mg, 2.2 Ommol) in foluene (16mL). The reaction mixture is degassed 5 times and flooded with N₂ again. Add bromide (30) (498mg, 2.00mmol) and finse with foluene (4.5 mL), degas again (5 times) and sitr the reaction solution 21h at 100°C. Partition the reaction solution between EIOAc and brine (1+1) and oxitact the separated squeeous layer 3 times with EIOAc. Wash combined organic layer with brine and dry with Na₂SO₄. Remove solvent under reduced pressure and purify the cruce product by preparative radial chromatography (allice get 60FC, CyHCIDAC 6+1, late 3+1) to obtain product (31) as a light yellow solid (300mg, 57%). [N. Miyaura; A. Suzuki; Chem. Rev.; 1995; 52457]. "In NMR (400MHz, CDCQ): 1.28 (1, 3 H, J = 7, 1 Hz); 6.77.6. At (m, 2 H); 6.91 (d, 1 H, J = 5, 1 Hz); 7.04 (d, 1 H, J = 5, 1 Hz); 7.61 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 5, 1 Hz); 7.61 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 5, 1 Hz); 7.61 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1 H, J = 6, 1 Hz); 7.04 (d, 1

J = 7.8 Hz, 13 Hz); 7.25 (d, 1 H, J = 7.8 Hz), 1903 [1934] Type (1.9 Hz) (1.9 H

M. J. F. T. Hz.); 6.58 (d, 1 H, J = 8.6 Hz); 6.59 (d, 1 H, J = 3.5 Hz); 6.75 (d, 1 H, J = 3.5 Hz); 6.56 (d, 1 H, J = 8.6 Hz); 7.05 (l, 1 H, J = 7.8 Hz); 7.26 (dd, 1 H, J = 7.8 Hz, 1.3 Hz); 7.30 (dd, 1 H, J = 7.8 Hz, 1.3 Hz); 7.90 (br.s.; 1 H), 8.38 (d, 1 H, J = 8.8 Hz).

[0035] Step 5: Dissolve ester (32) (118 mg, 0.25 mmol) in methanol (8.0 mL), add a 1M aqu. LiOH solution (1.75 mL, 1.75 mmol) and stir 20h at it. Remove solvent under reduced pressure und partition residue between CHCl₃ and 0.5M HCl (1+1). Separate the aqueous layer and extract 3 times with CHCl₃. Wash the combined organic layer with brine and dry with Na₂SO₄. Remove solvent under reduced pressure and dry the residue without further purification in oil pump vacuum to obtain crude product (33) as light brown foam (120 mg, quant). ¹H hMR (400MHz, COC)₃ :58 (s, 2 H); 3.78 (s, 3 H); 3.78 (s, 2 H); 3.86 (s, 3 H); 5.86 (s, 1 H); 5.76 (d, 1 H, J = 8.3 H2); 6.97 (d, 1 H, J = 3.5 H2); 7.22-7.27 (m, 1 H); 7.31 (td, 1 H, J = 7.8 H2, 1.3 H2); 7.86 (f, 1 H, J = 8.9 H2). H); 8.37 (d, 1 H, J = 8.9 H2).

[0036] Step 6: (The following reaction is done in an anhydrous N₂ atmosphere.) Dissolve (33) (118mg, 0.27mmol) anhydrous dichioromethane (2.5mL), cool to -78°C (acatione dry ice) and add slowly a 1mt BBrg solution in dichloromethane (1.61mL, 1.61mol). Stir the reaction mixture for additional 30min at -78°C. Remove cooling bath and stir reaction mixture to 6°C, and stowly water (1.00mL) under vigorous stirring. Partition the reaction mixture between EIOAc and water (1+1). Extract the separated squeous fayer with EIOAc (2 times) and wash the combined organic layer with brine and dry with Nay-SC, Remove solvent under reduced pressure and purity roude product by preparative RF HPLC (gradient, water/CH₂CN) 95.5) to obtain 6-[2-[2-(2.24, 3-firrilyd oxypheny)-acetylaminol-phenyl thiophen-2-yll acetic acid (34) as a light brown toam (Samg, 50%). H NMR (400MHz, CDCI₃): 3.54 (c, 2 H), 6.34 (d, 1 H, J = 8.3 Hz), 6.54 (d, 1 H, J = 8.3 Hz); 6.75 (d, 1 H, J = 3.3 Hz); 7.18 (d, 1 H, J = 7.5 Hz); 7.32 (t, 1 H, J = 7.8 Hz); 7.39 (d, 1 H, J = 7.3 Hz); 7.90 (d, 1 H, J = 8.1 Hz); 6.75 (c) s. 1 Hz).

4-(3.4.5-Trihydroxy-benzovlamino)-butyric acid (36)

[0037]

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SCHEME 8 HCPH₂N 111 0 El₃N 0H 35 UOH 0H 36

[0038] Step 1: (The following reaction is done in an anhydrous N₂ atmosphere.) Dissolve 4-amino-butyric acid ethyl ester hydrochioride (11) (3.98g, 23.74mmol) in anhydrous dichloromethane (120mL) at rt and add triethylamine (8.30mL, 59.55mmol). Sit rthe reaction solution for 15min at rt, add portionwise 3,4,5-trimethoxy-benzoyl chloride (6.58g, 28.53mmol) and stir for additional 24h at rt.

Hydrolize the reaction solution with methanol, filtrated the mixture through a short pad of silica gel with EIOAc/CyH (3+1). Purify crude product by flash chromatography (silica gel, EIOAc/CyH 2+1, later 3+1) to obtain (35) as a white solic (4.48g, 55%). H NMR (400MHz, CDCl₂), 1.22 (1, 3 H, J_0 = 7.1Hz); 1.95 (Hquim, 2 H, J_0 = 6.5Hz), 2.44 (t, 2 H, J_0 = 6.5Hz), 3.85 (t, 2 H, J_0 = 6.5Hz), 3.85 (t, 3 H); 3.89 (t, 6 H); 4.10 (q, 2 H, J_0 = 7.1Hz); 6.72 (br.s, 1 H); 7.02 (s, 2 H).

[0039] Step 2: Dissolve (35) (4.48g, 13.79mmol) in acetonitile (100mL) at rt and add 1M aqu. LiOH (41.4mL, 41.4mmol). Stir reaction mixture for 18h at rt.

Hydrolize the reaction mixture (cooling bath) with 1M siqu. HCI (50mL). Extract the mixture with chiproform (3 times), wash the combined organic layers with britine and dry with Na₂SQ, to obtain 4/3,4.5-intyfoxy-benzoylamino)-butync acid (36) as a white solid (4 07g, 979-5). No further purification. ¹H NMR (400MHz, CDCly): 1.94 (Yquint, 2H, J= 6 7Hz): 24 61, 12 H, 2 6 7Hz): 35 (0 ftrd, 2 H): 35 (6; 3 H): 38 (6; 6, H): 6.61 (No. 5, 1 H): 7.00 (6; 2 H).

3-[4-(3,4,5-Trihydroxy-benzoylamino)-butyrylamino]-benzoic acid (38)

[0040]

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[0041] Step 1: (The following reaction is done in an anhydrous N₂, atmosphere.) Dissolver EDC x Hcl (141mg, 0.74mmol) and El₃N (0.105mL, 0.74mmol) in anhydrous DCM (4.0 mL) and stir for 5min at rt. Added (36) (208mg, 0.70mmol) and DMAP (8 8mg, 0.07mmol) and stir for 5min Add 3-Aminomethyl-benzoate (159mg, 1.05mmol) and stir the reaction solution for 21h at rt. Hydroite the reaction solution with sat. aqu NH₂Cl and water, separate layers, extract and layer with EICA c2 (times) and water in combined organic layer with vater and brine and dry with N₃S₂O₂, Purified crude product by flash chromatography (silica gel 60, EICAC/CyH 3-1, later EICAC/MeOH 9-1) to obtain G7) as a white solid (279mg, 7.2%). If NMH (400MHz, CCCl₃); 2.01-2.10 (br.m, 2.81), 2.52 (br.1, 2.41, 2.6. e1cl₃); 3.53-3.63 (br. m, 2.11); 3.85 (s, 6.11); 3.89 (s, 3.11); 7.02 (s, 2.11); 7.35 (dd, 1.11, J₂ = 8.1142, J₂ = 7.612); 7.75 (d, 1.11, J=7.612); 7.80 (d, 1.11, J=1.1142, J₃ = 7.612); 7.80 (d, 1.1142, J₃ = 7.612)

[0042] Step 2: (The following reaction is done in an anhydrous N₂ atmosphere.) Dissolve (37) (86mg, 0.2mmol) in anhydrous DCM (4.0mL) at :25°C and add dropwise BBr₃ (0.303mL, 3.20mmol). The reaction mixture is slowly warmed up to 1 and stir for additional 6h at rt. Cool reaction mixture to 0°C, add dropwise water/ThF (1+1, mL) followed by DCM (2mL). Sit overnight and add MoOH (2mL) to homogenize the mixture. Remove solvent under reduced pressure and purity crude product by repearative RP HPLC (gradient, water/MeON 95.5) to obtain 3/4/(3.4.5-Tirhydroxy-benzoylamino)-buryylamino]-benzoic acid (38) as a white solid (41mg, 55%). 14 NNR (400MHz, CD₂OD): 2.02 (br.tt, 2 H, J₁ = 7.1Hz), 2.6. 6.5Hz); 6.90 (s, 2 H); 7.43 (dd, 1 H, J₇ = 8Hz), 7.86 (d.1, H, J, ca. 6.5Hz); 6.90 (s, 2 H); 7.43 (dd, 1 H, J₇ = 8Hz), 7.86 (d.1, H, J₈ = 8Hz), 7.86 (d.1, H, J₈ = 8Hz), 7.86 (d.1, H, J₈ = 8Hz); 8.26 (br.s. 1 H).

4-{[4-(2,3,4-Trihydroxy-benzoylamino]-butyrylamino]-methyl}-cyclohexanecarboxylic acid (46)

[0043]

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SCHEME 10

[0044] Step 1: Dissolve 4-Aminomethyl-cyclohexanecarboxylic acid ([39]; 1.15g, 7.30mmol) in 1.4-dioxane (12 om.) and 10% agu, Na₂Co₂ (23 cm.) and add Fmo-Ci (2.25g, 8.76mmol). Sit the reaction mixture for 2.5 h at 1. Add 1 M agu HCI (42 0ml,) to the mixture and extract with E10Ac (3 times). Wash the combined organic layers with 1 M agu HCI (42 0ml,) to the mixture and extract with E10Ac (3 times). Wash the combined organic layers with 1 M agu HCI, water and brine, extract the combined and layers once again with E10Ac, dry the combined organic layers with Na₂SO₄ and remove solvent under reduced pressure. The left crude product is washed with ice cold E10Ac and dried in oil pump vacuum to obtain (40) as a white sold (2.25g, 83°). No further purification, IM, Nichtofre, E1. Schacht, Teirahetron; 1994; 50: 12; 3747-3760; 14 NMRI (400MHz, CDCl₃): 087-1 cg (br. m, 2 H); 1.34-1.52 (br. m, 4 H); 1.55 (br. m, 4 H); 1.56 (br. m, 4 H); 1.57 (br. m, 4 Hz); 7.57 (d, 2 H, J = 7.4Hz); 7.57 (d, 2 H, J = 7.4H

[0045] Step 2: Dissolve (40) (190mg, 0.5mmol) in DCM (0.5mt) and DMF (0.25mt) at r in a pre-dried tube, add DIEA (0.52mt, 1.5mmol) and add this solution to 2-chlorotrilychlorid-polystyrene (82mg, 0.13mmol) which is preswollen before in DCM (0.15mt). Shake the reaction suspensions for 14h at rt. The resin is (41) washed with DCMMeOH/DIEA (17+2+1, 3 limes). DCM (once), DMF (3 limes) and again DCM (twice) and dried in vacuum. [i.e. Novabiochem/9 2000 Catalog, 2000; \$15-\$15.

[0046] Step 3: Suspende complete amount of resin (41) from step 2 in DMF (0.4mL) and piperidine (0.1mL) and shake it for 1.5h at n. Washed resin (42) with DMF (3 times) and DCM (3 times for 30min) and dry in vacuum

[0047] Step 4: Complete amount of resin (42) from step 3 was preswollen in DMF (0.8mL) for 20min. Dissolve acid (43) (175mg, 0.59mmol) and HOBI (90mg, 0.59mmol) in DMF (2.8mL) at rt, add DIC (75mg, 0.59mmol), stir for 15min and add this solution to the preswollen resin (42). Shake the resin suspension gently for 20h at rt. Wash resin (44) with DMF (3 times) and DCM 4 it limes) and dry in vacuum.

[0049] Step 5: (The following reaction is done in an anhydrous N₂ atmosphere.) Suspende complete amount of crude (45) from step 3 in anhydrous DCM (2.0mL), cooled it to -40°C and added BBr₃ (0.15mL, 1.59mmol). Shake the reaction suspension to 1 ha -40°C, 2 ha - 25°C and 30min at +50°C. Add dropwise water under vigorous stirring followed by MeOH. Remove solvent under reduced pressure. Purify the crude product by preparative RT:-(91C (3000)) 100 (40°C, 2 H. J. = 3.0Hz, J. = 12.9Hz); 1.30-1.55 (m, 3 H); 1.86 (br.d., 2 H. J. = 13Hz); 1.94 (*Yquint, 2 H. J. = 7.1Hz); 2.03 (br.d., 2 H. J. = 13Hz), 2.24 (*YPH, 1 H. J. + 3.3Hz, J. = 12.9Hz); 2.30 (h. H. J. = 8.9Hz). 7.16 (at 1.1 H. J. = 8.9Hz), 7.16 (at 1.1 H. J. = 8.9Hz). 7.16 (at 1.1 Hz). 7.16 (at 1.1

Sialyl Lewis Tyrosine Sulfate Assay (sLc TSA):

[0050] Compounds of the present invention are assayed on a molecular level for their ability to inhibit the binding of P. L., or E-selectin chimeric molecules to site and tyrosinesuffate residues linked to a polymeric matrix as a PSGL-1 substitute. Selected 50% inhibitory concentrations (C₅₀-yrailues) are determined.

Microtiler plates are content overnight in carbonate butfer pH9.6 with goal and human Fc mAB (10 µg/m), After washing in assay butfer (25mM 4-(2-hydroxyelthyl)-1-pipera/ineathanesulfone and (HePES), 15mM NoC1, mM CaCl, pH7.4) and blocking (3% bovine serum albumin (6SA) in assay butfer) plates are incubated for 2h at 37°C with human P-Selectin-IgG-chimera (0.6 In Mr espectively 150ng/mL) or human L-Selectin-IgG-chimera (0.6 In Mr espectively 180ng/mL) or human L-Selectin-IgG-chimera

[0051] The compounds referred to in the following SCHEME11 are those compounds referred to as the particularly preferred compounds herein.

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SCHEME 11

| Continue | Continue

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Compound	E-Selectin [% inhib.]	STD	P-Selectin [% Inhib.]	STD	L-Selectin [% inhib.]	STD
34	46.5	4.4	92.4	0.2	81.9	0.6
47	56.2	9.4	97.4	0.4	87.7	4.2
48	79.4	1.7	78.2	2.7	76.2	1.6
49	54.4	10.0	84.9	1.4	88.4	2.5
50	89.2	0.4	90.5	1.0	81.7	0.5
51	63.1	9.4	86.2	2.2	73.7	2.8
52	32.9	7.0	87.4	0.2	81.0	1.6
53	58.5	1.4	,57.7	4.3	52.3	3.8
54	58.9	4.3	96.5	1.0	83.6	1.0
55	37.6	4.1	51.2	1.3	48.7	7.0
56	50.4	1.8	93,1	0.9	95.2	0.5
57	99.8	2.1	97.9	0.4	96.5	0.2
58	68.9	5.0	87.3	0.7	81.7	0.9
59	67.5	2.3	67.1	8.4	62.4	3.4
60	90.2	4.0	86.3	2.2	84.9	1.9
61	52	0.6	94.6	2.5	85.4	4.7
62	53.4	7.6	87.2	0.7	75.4	1.0
63	85.4	5.3	86.6	1.6	84	2.2
64	82.9	1.4	73.1	1.4	72.6	2.8
65	96.4	3.3	96:1	0.6	92	1.0
66	46	6.9	99.1	0.1	97.4	0.3
67	38.7	4.0	97.7	0.2	83.8	1.2
68	42.2	5.6	73.8	3.1	48.8	3.1
69	38.7	9.1	99.2	0.1	95.0	0.4
70	n.a.		89.4	2.1	88.7	3.6
71	92.7	2.8	88.6	1.9	87.4	1.5
72	94.9	5.1	86.6	3.7	89.9	3.9
73	53.5	0.7	79.4	2.8	66.2	3.2

Compound	IC ₅₀ E-Selectin [μM]	STD	STD IC ₅₀ P-Selectin [µM] ST		IC ₅₀ L-Selectin [μM]	STD
34	n.a.		7.7	1.0	11.6	3.3
51	0.8	0.2	1,1	0.0	1.4	0.3
53	1.0	0.7	2.1	0.6	n.a.	
57	3.3	2.0	2.4	0.7	3.1	1.0
59	3.0	1.2	2.8	1.1	3.3	0.9

(continued)

Compound	IC ₅₀ E-Selectin [μM]	STD	IC ₅₀ P-Selectin [µM]	STD	IC ₅₀ L-Selectin [µM]	STD
60	3.1	2.4	3.4	2.1	5.7	1.0
63	1.3	0.3	1.7	0.5	1.8	0.2
64	2.3	1.0	2.3	0.6	2.3	0.7
66	0.6	0.2	2.1	0.7	7.0	7.7
70	n.a.		7.4	1.0	16.0	4.8
71	7.3	9.4	13.0	17.2	15.0	20.0
72	8.4	5.6	21.7	25.7	109.4	198
73	12.2	16.0	14.8	13.9	26.2	32.5

Flow Chamber Assay / Cell Adhesion and Rolling under Flow Conditions

[0052] To assess the capability of compounds to inhibit cell binding under dynamic conditions resembling the flow in a blood vessel, flow chamber assays addressing testing binding of HL-60 cells / various cell times to P-selectin, L-selectin and E-selectin and E-selectin charge in performed.

Cell attachment under flow conditions are determined using a parallel flow chamber system. A 35mm polystyrem culture dish is coarde for 1 hour at nom temperature with coaling butter (50mM tris-flydroxymethyl) aminomethane buffer (Tris), 150 mM NaCl, 2 mM CaCl₂; pH 7,4) containing human E- or P-selectin-tgG chimera at concentrations of 2,5,gmlm or 1)gmlm, respectively, Atter removal of the coaling solution non specific binding sites are blocked for an additional hour with 15,6 BSA in coaling buffer at room temperature. After washing with assay buffer (*Roswell Park Memorial Institute 1640", (RPM 1640) + 10mM HEPES) the dish is fitted into a parallel plate learning flow champer (sold from Glycotich, Rockville, MD) and mounted on an inverted phase-contrast microscope (sold from Olympus, Hamburg, Germany) equipped with a CCD camera (JVC) that is connected to a PC. Employing a peristalitic pump (35gM compound or vehicle control (DMSQ). Cells (1 millior) m1) are added to the chamber and allowed to distribute for 2 minutes at a high flow rate. The flow rate is then decreased resulting in a calculated flow shear of 1 dymecrne. Video sequences of 10 low power fields are digitally recorded after 5 minutes continuous flow. The precentage of inhibition is calculated from the mean number of cells per field that attached to the coated dish surface in the presence versus absence of compound of at independent experiments.

Data from Flor	w Chamber Assay fo	r E- and P-Selectin
Compound	E-Selectin [% inhib.]	P-Selectin [% inhib.]
57	35	11 '
63 .	n.a.	17
71	22	22

Claims

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Compounds having the general structure of formula (I).

wherein the symbols, indices and substituents have the following meaning if R^2 =OH and R^3 =H then R^1 =H, CN, NO₂, CF₃, F, Cl, Br, I, CH₃ or

if R3=OH and R3=H then R3=H, CN, NO₂, CF₃, F, Cl, Br, I, CH₃, Et, n-Pr, i-Pr, n-Bu, i-Bu, t-Bu, phenyl, thienyl, furyl, thiazolyl or

if R³=OH and R¹=H then R²=H, CN, NO₂, CF₃, F, CI, Br, I, CH₃, Et, n-Pr, i-Pr, n-Bu, t-Bu, phenyl, thienyl, furyl, thiazolyl

(a)

with -E- being -NH-, - $(CH_2^-)_k$ NH-; -G- being - $(NH-)_m$ and g = 0,1; h = 1, 2, 3; k = 1, 2, 3; m = 0,1; n = an integer from 1 to 8

(b)

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with R4 being H, CH3, CH2CH3

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z-KI-GI-

with $\rm H^5$ being H, NO₂, CF₃, F, CI, Br, I, CN, CH $_{\rm 3}$ and -K- being -S- or -O-

(d)

with p being an integer from 2 to 8

(e)

$$r^{r_{z}}(E) \int_{0}^{g} \left(\int_{0}^{g} \int_{0}^{g} \left(\int_{0}^{g} \int_{0}$$

with q being an integer from 1 to 9 and r being an integer from 1 to 3 $\,$

(f)

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$$\mathcal{F}_{\mathcal{F}_{k}}(\mathbb{E}_{0}^{\mathbb{F}_{k}}(\mathbb{F}_{1}$$

with -T1-, -T2-being -E-, -K- or -(N-alkyl)-

with -V- being -(NH-)s- with s being 0 or 1,

R[§] being Co₂H, CO₂Miyl, CO₂Anyl, CO₂Mn₂, CO₂Anallyl, So₃H, So₂Mh₂, PO(OH)₂, 1-H-tetrazolyl-, CHO, COCH₃, CH₂OH, SH, NH₂, NHAllayl, N(Allayl)Mayr, OcH₃, CH₂OH₃, SH R⁷ independently from R⁸ being H. CH₃, CH₂CH₃, CF₃, F, Cl, Br, I, CN, NO₂ and R⁸ independently from R⁸ and R⁹ being H. CH₃, CH₂CH₃, CF₃, F, Cl, Br, I, CN, NO₂, R⁶ R⁹ being H, NO₂, CF₃, F, Cl, Br, I, CN, CH₃, OCH, SH I being 0.1.2 and ·W- = (CH₂-)₂, cis-CH=CH- or trans-CH=CH-, and v being 0.1.2

-Z =

$$\text{H}_{\text{B}_{0}} \text{M}_{\text{-B}_{0}}$$

-W-R

and the pharmaceutically acceptable salts, esters or amides and prodrugs of the above identified compounds of formula (f).

2. Compounds according to claim 1, wherein the compounds are defined by formulas (A) and (B) $30\,$

wherein -Y is (Q=CH, N)

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with R10 being CO₂H, CO₂aikyl, CO₂aryl, CO₂NH₂, CO₂aralkyl, CH₂SO₃H, CH₂SO₃NH₂, CH₂PO(OH)₂, 1-H-letrazolyl, CHO, COCH₃, CH₂OH, CH₂OH, CH₃MHalkyl, CH₂Mklkyl)aikyl', CH₂OCH₃, CH₂SH, R11 being CO₂H, CO₂alkyl, CO₂aryl, CO₂NH₂, CO₂aralkyl, SO₃H, SO₂NH₂, PO(OH)₂, 1-H-letrazolyl, CHO, COCH₃, OH, NH₃, NHalkyl, Nalkylyaikyl', OCH₃, SH.

3. Compounds according to claim 1, wherein the compounds are defined by formulas (C) and (D)

wherein Y is like relating to formulas (A) and (B) and X is:

- Pharmaceutical compositions comprising at least one compound of the formala (I) and a pharmaceutically acceptable carrier which is useful in a medicine.
- Pharmaceutical compositions according to claim 4, comprising at least one compound of formula (A) or formula
 (B) or formula (C) or formula (D).
 - Method of inhibiting the binding of P-selectin, L-selectin or E-selectin to sLe* or sLe* and tyrosinesulfate residues
 comprising the step of administering to a patient an effective amount of at least one compound having the structure
 of formula (i).
- Use of compounds according to any one of claims 1 to 3 for the preparation of a medicine for the treatment of a
 patient, inhibiting the binding of P-selectin, L-selectin or E-selectin to sLe^x or sle^a and tyrosinesulfate residues.

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European Patent

PARTIAL EUROPEAN SEARCH REPORT

Application Number

which under Rule 45 of the European Patent Convention EP 04 00 6461 shall be considered, for the purposes of subsequent proceedings, as the European search report

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Υ	*columns 1 and 2; with rn 214124-47-	the claims and compound	2,3	070237700
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INCOL	MPLETE SEARCH			
be carried	th Division considers that the present y with the EPC to such an extent that out, or can only be carried out partix prohed completely:	application, or one or more of its olaims, does/d a meaningful search into the state of the ort con thy, for these claims.	lo not	
Claims se	maked incompletely:			
Claims no	searched :			
Reason fo	the Emitation of the search:			
see	sheet C			
			j	
	Place of search	Date of completion of the sounds		Example
	Munich	15 September 2004	Lore	enzo Varela, M.J.
CA	TEGORY OF CITED DOCUMENTS	T theory or principle :	underlying the in	vention
X : partie	ularly relevant if taken alone	E: earlier patent docu after the täing date		hed an, or
Y : partic	ularly relevant a combined with anot ment of the same category	her D; document cited in t	the application	
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		document		



European Pater

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Application Number EP 04 00 6461

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PARTIAL EUROPEAN SEARCH REPORT

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PARTIAL EUROPEAN SEARCH REPORT

Application Number

Office EP 04 00 6461

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INCOMPLETE SEARCH SHEET C

Application Number EP 04 00 6461

Although claim 6 is directed to a method of treatment of the human/animal body (Article 52(4) EPC), the search has been carried out and based on the alleged effects of the compound/composition in the same extent as explained herein below for the compound claims 1-3.

Claim(s) searched incompletely:

Reason for the limitation of the search:

Present claims 1-7 relate to an extremely large number of possible compounds/compositions and their use in medicine. Support within the meaning of Article 84 EPC and disclosure within the meaning of Article 83 EPC is to be found, however, for only a very small proportion of the compounds claimed (those ones in the activity examples). In the present case, the claims so lack support, and the application so lack disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, the search has been carried out for those parts of the claims which appear to be supported and disclosed, namely those parts relating to the compounds disclosed in the activity examples). In fact, the claims contain so many options, variables, possible postate that a dack error possibilities of attachment of the different arises to such an extent as to render the meaning of Article 84 EPC arises to such an extent as to render the meaning of Article 84 EPC arises to such an extent as to render the meaning of Article 84 EPC arises to such an extent as to render the above, the search has beliams impossible. Consequently, as explained above, the search has beliams out for those parts of the application which do appear to be clear namely the compounds in the activity examples and closely related homologous compounds; compounds according to formulae A, B wherein Y is as in claim 2 with R10 and R11 being CQ20 or CQ201kyl.

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 04 00 6461

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